BABYNET Covered Diagnoses

Children with documentation that any condition on this list has been professionally diagnosed are eligible for BabyNet services based on "established risk".

10p13 Deletion Dandy walker malformation 11q Deletion Down syndrome (trisomy 21)

13q Syndrome Duplication short arm chromosome #20

18q Deletion Syndrome Encephalocele 49xxxxy syndrome Fazio~Londe disease Fetal alcohol syndrome 4p Minus Syndrome

6q Minus Syndrome Fragile X

7q Minus Syndrome

Agenesis of the corpus callosum Grade IV intraventricular hemorrhage (IVH)

Glaucoma w/visual impairment

Albinism Hearing loss \geq 26 db

Amniotic band syndrome Hemiparesis

Amyoplasia congenita disrutice sequence Herpes encephalitis Anencephaly Holoprosencephaly

Angelman syndrome Hydranencephaly Anophthalmia Hydrocephaly

Arginosuccinic aciduria Incentinentia pigmenti syndrome

Arthrogryposis Infantile spasms Asphyxia Isochrome 18 p syndrome Athetoid CP Kabuki syndrome

Auditory neuropathy

Karsch-neugebauher syndrome Autism spectrum disorders (ASD)

Klinefelter syndrome Automatic eligibility nos Krabbe's disease Bilateral micromelia Larsen syndrome Bilateral optic nerve coloboma Lebers amaurosis

Bilateral retinal detachment w/blindness Lennox-gastaut syndrome

Bilateral visual acuity ≤ 20/70 corrected vision best eye Lissencephaly syndrome

Lowe syndrome (oculo-cerebro-renal) Birthweight ≤ 1200 grams Marshal Smith Syndrome Carpenter's syndrome

Melnick-Frazier Cataracts w/ visual impairment Microdactvlv Caudal regression syndrome Midas syndrome Cerebral palsy (CP)/static encephalopathy

Miller-dieker syndrome Charge association

Mobius sequence Citrulinemia Mosaic trisomy 8 Cleft hands bilertal

MPS (mucopolysaccharidosis) Coffin Lowry syndrome MSUD (maple syrup urine disease) Cornelia de Lange

Myelodysplasia Cortical blindness Myotonic dystrophy Cri du chat Myotubular myopathy Cystinosis

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Neural tube defects

Opitz syndrome

Optic nerve atrophy

Ornithine-carbamyl-transferase deficiency

Osteogenesis imperfecta

Pachygyria

Pallister-killian syndrome

Pathologic head growth

Perinatal asphyxia, severe

Pervasive developmental disorder (ASD)

Phocomelia

PKU

Prader-willi syndrome

Prematurity (gestational age \leq 28 wks)

Propionic acidemia

R.O.P. stage 5 & 6 retrolental fibroplasia

Retinitis pigmentosa

Retinoblastoma

Rhizomelic chondrodysplasia punctata

Ring chromosome 13

Schizencephaly

Seckel syndrome

Seizures w/congenital brain malformation

Septo-optic dysplasia

Severe attachment disorder (ASD)

Spastic diplegia

Spastic hemiplegia

Spastic quadriplegia

Spina bifida

Spinal cord injury

Spinal muscular atrophy

Stickler syndrome

Syringohydromyelia

Tar syndrome

Tay- sachs disease

Tetrasomy 12p

Trisomy 1

Trisomy 10

Trisomy 13

Trisomy 18

Trisomy 4

Trisomy 8

Trisomy 9

Tuberous sclerosis

Turner's syndrome

Vater syndrome, with limb anomalies

Velo-cardio-facial syndrome

Waardenberg syndrome

Werdnig-hoffman

William's syndrome

Wolfe-Hershorn syndrome